



Genomic Composition And Dynamics Among Methanomicrobiales Predict Adaptation To Contrasting Environments

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Abstract

Members of the order Methanomicrobiales are abundant, and sometimes dominant, hydrogenotrophic (H₂-CO₂ utilizing) methanoarchaea in a broad range of anoxic habitats. Despite their key roles in greenhouse gas emissions and waste conversion to methane, little is known about the physiological and genomic bases for their widespread distribution and abundance. In this study, we compared the genomes of nine diverse Methanomicrobiales strains, examined their pangenomes, reconstructed gene flow and identified genes putatively mediating their success across different habitats. Most strains slowly increased gene content whereas one, *Methanocorpusculum labreanum*, evidenced genome downsizing. Peat-dwelling Methanomicrobiales showed adaptations centered on improved transport of scarce inorganic nutrients and likely use H⁺ rather than Na⁺ transmembrane chemiosmotic gradients during energy conservation. In contrast, other Methanomicrobiales show the potential to concurrently use Na⁺ and H⁺ chemiosmotic gradients. Analyses also revealed that the Methanomicrobiales lack a canonical electron bifurcation system (MvhABGD) known to produce low potential electrons in other orders of hydrogenotrophic methanogens. Additional putative differences in anabolic metabolism suggest that the dynamics of interspecies electron transfer from Methanomicrobiales syntrophic partners can also differ considerably. Altogether, these findings suggest profound differences in electron trafficking in the Methanomicrobiales compared with other hydrogenotrophs, and warrant further functional evaluations.

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Genomic composition and dynamics among *Methanomicrobiales* predict adaptation to contrasting environments

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Members of the order *Methanomicrobiales* are abundant, and sometimes dominant, hydrogenotrophic (H₂-CO₂ utilizing) methanoarchaea in a broad range of anoxic habitats. Despite their key roles in greenhouse gas emissions and waste conversion to methane, little is known about the physiological and genomic bases for their widespread distribution and abundance. In this study, we compared the genomes of nine diverse *Methanomicrobiales* strains, examined their pangenomes, reconstructed gene flow and identified genes putatively mediating their success across different habitats. Most strains slowly increased gene content whereas one, *Methanocorpusculum labreanum*, evidenced genome downsizing. Peat-dwelling *Methanomicrobiales* showed adaptations centered on improved transport of scarce inorganic nutrients and likely use H⁺ rather than Na⁺ transmembrane chemiosmotic gradients during energy conservation. In contrast, other *Methanomicrobiales* show the potential to concurrently use Na⁺ and H⁺ chemiosmotic gradients. Analyses also revealed that the *Methanomicrobiales* lack a canonical electron bifurcation system (MvhABGD) known to produce low potential electrons in other orders of hydrogenotrophic methanogens. Additional putative differences in anabolic metabolism suggest that the dynamics of interspecies electron transfer from *Methanomicrobiales* syntrophic partners can also differ considerably. Altogether, these findings suggest profound differences in electron trafficking in the *Methanomicrobiales* compared with other hydrogenotrophs, and warrant further functional evaluations.

Introduction

Methanogens consist of at least nine (Mihajlovski *et al.*, 2008; Sakai *et al.*, 2008; Evans *et al.*, 2015; Nobu *et al.*, 2016) recognized orders and are divided into two major categories based on whether or not they contain cytochromes. Methanogens of the *Methanosarcinales* order contain cytochromes and use substrates such as acetate, methanol,

methylamines and H₂/CO₂. *Methanosarcinales* have been extensively studied for their biochemistry and genetics (Ferry, 2011). All other orders of methanogens lack cytochromes and typically reduce CO₂ using H₂, formate and secondary alcohols (Thauer *et al.*, 2008), although some are capable of reducing methylated substrates with H₂ (Fricke *et al.*, 2006; Brugère *et al.*, 2014). Much of what is known about cytochrome-deficient methanogens comes from biochemical and genetic experiments performed on members of the *Methanobacteriales* and *Methanococcales* orders (Ferry, 2011), compared with which the cytochrome-deficient *Methanomicrobiales* are highly understudied.

Comparative genomic analysis across a microbial group (that is, pangenome analysis) has been proven valuable to identify functions that over time have led

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to niche specialization at broad (Luo *et al.*, 2013) or fine (Lee and Marx, 2012; Youngblut *et al.*, 2015) phylogenetic levels. Genome expansion can lead to metabolic versatility, or the ability to take advantage of high nutrient availability, or other adaptations not directly related to metabolism (Ochman *et al.*, 2000; Dobrindt *et al.*, 2004), while genome downsizing in free-living microbes can cause specialization by natural selection eliminating unnecessary genes during colonization or residency in an environment (Lee and Marx, 2012). Comparative genomics studies within different orders of methanogens (*Methanosarcinales*, *Methanocellales* and *Methanomassiliicoccales*) have shown major differences in genome content between members, and evidenced metabolic adaptation to low substrate (H_2) concentrations, evidenced novel energy conservation pathways and evidenced genomic adaptations to different environments (Maeder *et al.*, 2006; Borrel *et al.*, 2014; Lyu and Lu, 2015; Youngblut *et al.*, 2015).

Methanogens of the order *Methanomicrobiales* are widespread across terrestrial, freshwater, marine and bioreactor ecosystems (Garcia *et al.*, 2006). Phylotypes associated with this order have been found highly abundant or dominant across very distinct conditions in anoxic soils of wetlands, lakes or river sediments, peatlands, waste-treating bioreactors, among others (Bräuer *et al.*, 2006; Auguet *et al.*, 2009; Bialek *et al.*, 2013). Although these methanogens have a narrow set of metabolic substrates (Garcia *et al.*, 2006), their broad distribution indicates its members have differentially adapted to contrasting conditions ranging from low to high: pH, nutrients, salinity, oxidative stress, temperature and others. How do diverse members of the order *Methanomicrobiales* succeed in such a wide range of conditions? Answers to this question are needed to better understand their activity and predict potential changes in their role in CH_4 atmospheric emissions from natural or engineered environments or for technological purposes (Purwantini *et al.*, 2014).

Although the study of *Methanomicrobiales* genomics has received little attention, one early comparative analysis of three genomes available then revealed that the *Methanomicrobiales* represent a distinct class (class II) with their own functional and phylogenetic differentiations among methanogens (Anderson *et al.*, 2009a). Functional adaptations are likely across the diverse *Methanomicrobiales* and such can be explored among multiple genome sequences of strains from different habitats that have recently become available (detailed in Table 1). In this work, we developed the comparative analyses of nine *Methanomicrobiales* genomes to identify the order's putative genomic adaptations to contrasting environments.

Materials and methods

Genomes, protein clustering and phylogenetic inference

All annotated genome sequences (protein and nucleotide) were obtained from NCBI under accession numbers detailed in Table 1. Protein sequences were compared reciprocally using blastp (Altschul *et al.*, 1990) with an *e*-value cutoff of $1e-10$ and filtered for alignments that spanned at least 70% the length of the longer of the two proteins. Protein sequences were grouped into homologous clusters based on negative log transformations of their *e*-values with a ceiling of 200 using MCL version 11-294 (van Dongen, 2000). Based on proml (Felsenstein, 1989) phylogenetic inferences, core protein clusters containing paralogs were further refined manually by splitting robust branching groupings where appropriate.

Phylogenetic inference

Phylogenetic inference was performed using three data sets: (i) small subunit (SSU) rRNA, (ii) large subunit rRNA and (iii) core predicted protein clusters. To reduce phylogenetic noise in the core

Table 1 *Methanomicrobiales* strains analyzed in this study

Strain	Habitat	pH	$t_{gen}(h)$	Mbp	% GC	Accession number(s)	Ref.
<i>Methanoregulaceae</i> family							
<i>Methanoregula boonei</i> 6A8	Acidic bog	5.1	44	2.54	54.5	CP000780	Bräuer <i>et al.</i> (2011)
<i>Methanoregula formicica</i> SMSP	Anaerobic sludge	7.4	24	2.82	56.2	CP003167	Yashiro <i>et al.</i> (2011)
<i>Methanosphaerula palustris</i> E1-9c	Rich fen	5.7	19	2.92	58.9	CP001338	Cadillo-Quiroz <i>et al.</i> (2009)
<i>Methanolinea tarda</i> NOBI-1	Sewage sludge	7.0	144	2.05	56.3	AGIY01000000	Imachi <i>et al.</i> (2008)
<i>Methanomicrobiaceae</i> family							
<i>Methanoculleus marisnigri</i> JR1	Marine sediment	6.4	NA	2.48	62.1	CP000562	Romesser <i>et al.</i> (1979)
<i>Methanolacinia petrolearia</i> SEBR 4847	Off shore oilfield	7.0	10	2.84	47.4	CP002117	Ollivier <i>et al.</i> (1997)
<i>Methanoplanus limicola</i> M3	Mud of saline swamp	NA	NA	3.20	42.0	AHKP01000000	Wildgruber <i>et al.</i> (1982)
<i>Methanospirillaceae</i> family							
<i>Methanospirillum hungatei</i> JF-1	Sewage sludge	~7.0	17	3.54	45.0	CP000254	Ferry <i>et al.</i> (1974)
<i>Methanocorpusculaceae</i> family							
<i>Methanocorpusculum labreanum</i> Z	Tar pit	7.0	12.5	1.80	50.0	CP000559	Zhao <i>et al.</i> (1989)

Abbreviations: $t_{gen}(h)$, generation time (hours); Mbp, genome size in Mbp; NA, not available. pH optima are for growth in pure culture.

protein clusters data set, we omitted all clusters that at any point of our analyses had more than one copy per genome and filtered out clusters whose corresponding nucleotide codon alignment showed evidence of substitution saturation (Xia and Lemey, 2009). The remaining 437 protein clusters (Supplementary Data 1) were aligned and concatenated for analyses.

Sequence and codon alignments were performed with Clustalw (Larkin *et al.*, 2007) and Pal2nal v14 (Suyama *et al.*, 2006), respectively. Gapped positions were removed with T-coffee (Notredame *et al.*, 2000). The nucleotide index of substitution saturation test was done with Dambe 5.3 (Xia *et al.*, 2003, 2013). Parameters for phylogenetic inference from nucleotide or protein alignments were estimated using jModelTest (Darriba *et al.*, 2012) and ProtTest3 (Darriba *et al.*, 2011), respectively. Phylogenetic inferences based on proml, protpars, dnaml and dnajpars were done using the Phylip package (Felsenstein, 1989). Mega5 (Tamura *et al.*, 2011) was used to perform neighbor-joining phylogenetic inference. The Bayesian Markov chain Monte Carlo method was implemented with MrBayes 3.2 (Ronquist *et al.*, 2012). MrBayes was run using two independent runs with a burn-in value of 25% until the standard deviations of split frequencies remained below 0.01.

Assessment for positive selection

The topology presented in Figure 1a was used in the assessment of all core orthologous protein clusters for positive selection. Codon alignments were assessed for positive selection on every branch with codeml from the PAML 4.6 package (Yang, 2007), using two maximum likelihood reconstructions of protein evolution: one with neutral evolution model (null hypothesis) and the other using a model specifying a proportion of sites subject to positive selection (alternative selection hypothesis). Comparisons of likelihood ratio test between the two models

were the criteria for detecting positive selection in loci as detailed elsewhere (Xu *et al.*, 2011). The Benjamini–Hochberg procedure (Benjamini and Hochberg, 1995) was applied to control the false positive detection rate to 1%.

Reconstruction of gene flow dynamics

Reconstruction of gene flow dynamics was performed using COUNT (Csurös, 2010). Three categories of gamma distributed parameters were used for optimizing the rate parameters and gain, loss and duplication rates were allowed to vary between lineages.

Identification of horizontal gene transfer candidates

Putative horizontal gene transfer (HGT) candidates were screened for using three contrasting methods based on (i) detecting phylogenetically atypical proteins, (ii) detecting biases in dinucleotide frequencies and (iii) detecting biases in codon usage. Phylogenetically atypical protein sequences were detected using DarkHorse v1.4 (Podell and Gaasterland, 2007), using varying parameters to detect HGT acquisitions of different ages. When using DarkHorse, three filter thresholds (2%, 5% and 10%) were used and the definition of self was varied between four hierarchical levels of classification: (i) strain/species, (ii) genus, (iii) family and (iv) order. A normalized lineage probability index cutoff of 0.38 was applied, which was adequately stringent to examine HGT only from outside the domain *Archaea*. When a protein had a normalized lineage probability index <0.38 in at least two of the three filter thresholds, it was flagged as an HGT candidate. The age of HGT acquisition was estimated using the most recent definition of self at which the protein was flagged as an HGT candidate. A complementary analysis exploring the possible HGT history of a membrane-bound heterodisulfide reductase (HdrED) in *Methanosphaerula palustris* used

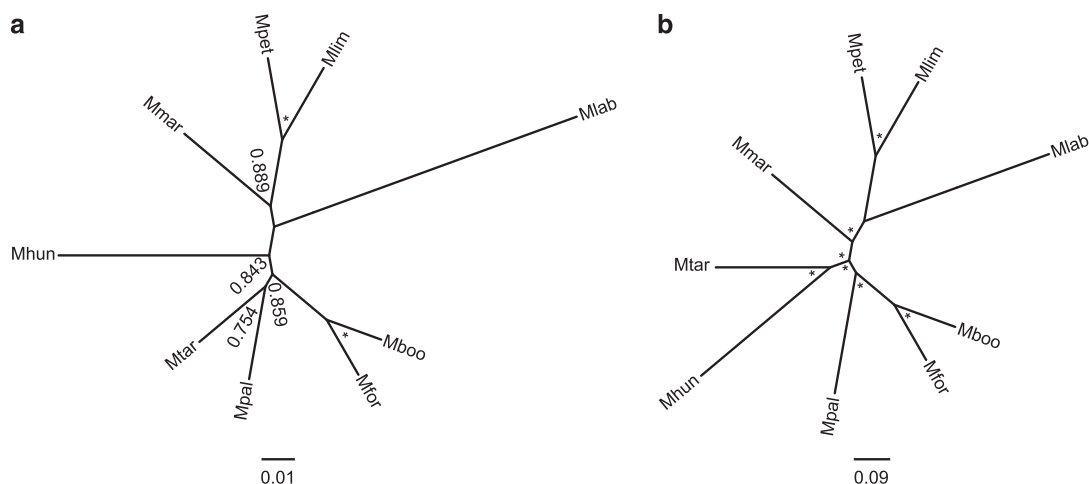


Figure 1 Phylogeny of the Methanomicrobiales. (a) Neighbor-joining (NJ) reconstruction of the small subunit rRNA gene. (b) Bayesian reconstruction of core genome amino-acid sequences. Bootstrap support for NJ reconstruction is shown next to nodes. Asterisks adjacent to nodes indicated either 100% bootstrap support or a calculated posterior probability of 1. Mboo, *M. boonei*; Mfor, *M. formicica*; Mpal, *M. palustris*; Mtar, *M. tarda*; Mmar, *M. marisnigri*; Mpet, *M. petrolearia*; Mlim, *M. limicola*; Mhun, *M. hungatei*; Mlab, *M. labreanum*.

a normalized lineage probability index cutoff of <0.85 . IslandPath-DIMOB was used to detect HGT related with the co-occurrence of abnormal dinucleotide frequencies and mobility genes (Hsiao *et al.*, 2003). SIGI-HMM as implemented in the Colombo package (Waack *et al.*, 2006) was used to detect HGT based on codon usage bias. For analysis of the *Methanolinea tarda* NOBI-1 draft genome assembly, only contigs greater than 100 kb in length were analyzed by IslandPath-DIMOB.

Inference of gene function

Functional annotations were parsed from the genbank format genome sequences. Functions assigned by KEGG orthology were used as annotations for the genomes in KEGG (Ogata *et al.*, 1999). Clusters of orthologous groups (COG) functional terms were parsed from the arCOG database (Wolf *et al.*, 2012). For genomes not in the arCOG database, the category assigned was the best hit of a blastp query against the arCOG database using an *e*-value cutoff of $1e-10$ and with at least 70% sequence overlap. Where necessary, the specificity of monovalent cation transporters was assessed using blast comparisons against experimentally verified transporters from bacteria.

Results

Pangenome and phylogenomics of the Methanomicrobiales

A total of 6749 predicted protein clusters were identified as part of the *Methanomicrobiales*' pangenome. Between 1012 and 1642 protein clusters were shared by each pairwise combination of strains (Supplementary Table 1). In all, 832 were common to all strains, representing 30–51% of their coding sequences, and are hereafter termed the *Methanomicrobiales*' protein coding core genome. In all, 699 of the core clusters were present as single orthologs in all strains; 2429 clusters were found in more than one but not in all strains, while from 244 to 532 clusters per strain were found to be unique to each of the nine strains (Supplementary Data 1). Rarefaction curves showed the core genome is well defined with the current data set, while the variable portions of the pangenome will increase with the addition of more strains (Supplementary Figure 1).

The phylogeny within the order was examined since evolutionary history is inextricably linked with genomic level adaptive processes. The strains had between a minimum of 85% and a maximum of 96% SSU RNA identity on a pairwise basis. Phylogenies constructed with several data sets (small or large subunit rRNA genes, and concatenated amino-acid sequences of 437 core orthologous clusters; Supplementary Data 1) and methods (detailed in Materials and methods) showed overall similar tree topologies with some incongruences (Figure 1; Supplementary Figure 2). The grouping of the *Methanoregulaceae* strains together is supported but the exact branching order of *M. palustris* and *M. tarda* is unclear. The placement

of *Methanoculleus* in the *Methanomicrobiaceae* family is uncertain, as *Methanoculleus marisnigri* did not always branch with *Methanoplanus limicola* and *Methanolacinia petrolearia*.

Gene flow dynamics in the Methanomicrobiales pangenome

The rates of gene acquisitions and losses between phylogenetic nodes (in Figure 1a) are presented in Figure 2. The *Methanomicrobiales* progenitor was inferred to possess 1744 protein clusters. Since the origin of the order, *Methanocorpusculum labreanum* has undergone a net loss of protein clusters while the other strains evidenced net gains, although some alternated between periods of net gain and net loss (Figure 2). Since the estimation of gene loss and gain rates relies on a robust phylogenetic inference, this analysis was repeated using all unique tree topologies inferred while assessing the order's phylogenetic structure (Supplementary Figure 2). The same main findings were consistently observed across iterations, although exact gene gain and loss rates varied to a minor extent depending on the tree topology used (for instance, 1725–1779 protein clusters were estimated for the progenitor ancestor).

This test was coupled with the screening for HGT candidates, which is a key for its role in encoding novel products into recipients' genomes. The three methodologies used in the HGT screening showed significant variation (Supplementary Figure 3). This reflects the fact that screening for phylogenetically atypical proteins can detect acquisitions of different ages while sequence composition biases may smooth out over time following acquisition (Lawrence and Ochman, 1997). The strains putatively acquired between 19.2% and 31.9% of their coding sequences via HGT, primarily from groups within the *Proteobacteria* and *Firmicutes* (Figure 3). Despite gene flow dynamics evidencing genome reduction in *M. labreanum*, this strain had the highest fraction of HGT candidates relative to its genome's size (Supplementary Figure 4). *M. labreanum* also showed a different profile of putative HGT donors, with gene gains from the *Firmicutes* representing a larger fraction than those from the *Proteobacteria* (Figure 3).

Assessment for positive selection in the protein coding core genome and differential content analysis in variable genome

Patterns of positive selection in the 699 core protein clusters present as single orthologs only were assessed on the various branches within the phylogeny presented in (Figure 1a). In all, 269 of the 699 core orthologous protein clusters had evidence of positive selection at some point with the false positive detection rate controlled to 1% (Supplementary Data 2). Of these, 106 were not localized to only one branch of Figure 1a. There were between 13 and 39 instances of protein

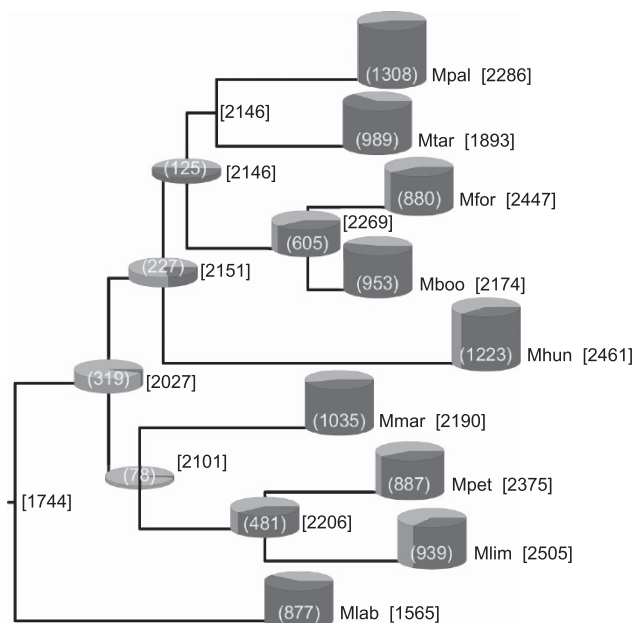


Figure 2 Reconstruction of the ancestral genome content of the *Methanomicrobiales*. Numbers in square brackets indicate the protein cluster content of the adjacent node or extant species. The number presented in parentheses represents the combined gene gains and losses inferred to have occurred since the previous node. Gains (black portion) and losses (white portions) are proportionally scaled in cylinders. The absence of a cylinder indicates that <0.5 gains plus losses were inferred to have occurred. Abbreviations defined in Figure 1.

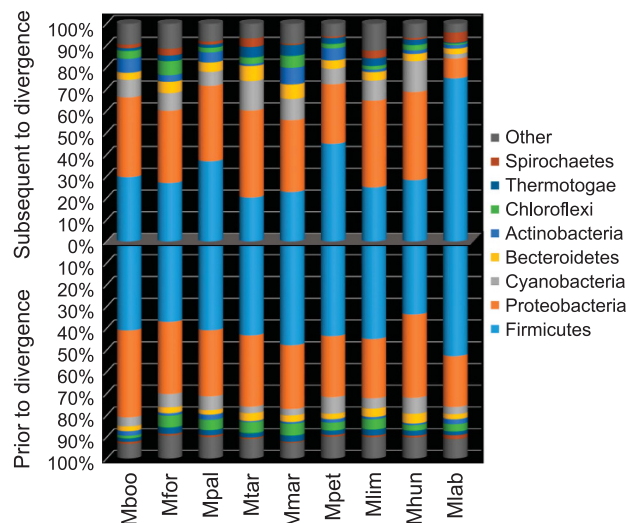


Figure 3 Phyla affiliations of most probable HGT donors. The phyla affiliations of putative bacterial HGT donors as inferred by DarkHorse to have occurred prior and subsequent to the divergence of the *Methanomicrobiales* order are presented. The numbers of HGT candidates are scaled to 100%. Abbreviations as in Figure 1.

clusters showing evidence of positive selection on only one of each of the individual branches.

The differential content analysis of non-core protein clusters (Supplementary Data 1) revealed 60 clusters were found only in both peat-dwelling strains, 13 were only found in strains tolerant of greater than 0.5 M NaCl and 11 clusters were present

only in strains isolated from bioreactors (Supplementary Table 2). Functional annotations of these genes suggested contrasting properties (Figure 4), although also a significant fraction did not present an annotation (57.4%) for better interpretation. The differential content analysis also revealed that *M. labreanum* lacks 96 protein clusters present in all other strains (Supplementary Table 3), while the other strains lacked only between 2 and 15 of such clusters. Notably, 14 of the 96 clusters absent in *M. labreanum* code for archaellum (archaeal flagellum) and chemotaxis proteins in the other order members, suggesting the loss of these functions.

We also analyzed the distribution of central metabolism genes including methanogenesis, energy conservation and electron trafficking (Figure 4, *Clusters 'S' in Supplementary Data 1*). Among these, three different membrane-bound hydrogenases, Ech (cluster S α), Eha (cluster S β) and Mbh (cluster S χ), were found. Only Ech was present in all strains (core) whereas Mbh and Eha were found only in six and five strains, respectively (Figures 4 and 5). A synteny analysis was performed on these membrane-bound hydrogenases and adjacent multisubunit Na⁺/H⁺ antiporters, if present, and on the HdrED genes that were all adjacent to genes encoding the D-subunit of the F₄₂₀-nonreducing hydrogenase (Mvh) (cluster S κ) (Figure 5). Furthermore, synteny analysis was also performed on the loci encoding MvhAG subunits (cluster S λ) (Figure 6) present only in two strains, *M. marisnigri* and *M. petrolearia* (Figure 4). The results of these analyses (discussed below) support hypotheses on the use of H⁺ rather than Na⁺-transmembrane chemiosmotic gradients, and the potential for differences in electron bifurcation steps in the central metabolism of members of this order.

Discussion

Pangenome and phylogenomics of the Methanomicrobiales

The *Methanomicrobiales* core' genome (832 genes) was found to be smaller than *Methanocellales* (1187 genes) (Lyu and Lu, 2015) but larger than *Methanomassiliicoccales* (658 genes) (Borrel *et al.*, 2014). However, an in-depth analysis of order-level core genome across methanogenic Archaea is limited by the uneven sampling of taxonomic diversity and number of genomes within groups, plus variation in criteria when delineating clusters. Meanwhile, inference from core genome content allowed assessing the phylogenetic structure of the strains in this study, which highlight some discrepancies previously noted elsewhere. For instance, analysis of SSU rRNA by neighbor-joining places *Methanospirillum hungatei* near but outside the *Methanoregulaceae* family while likelihood-based analyses place *M. hungatei* within such family (Cadillo-Quiroz *et al.*, 2009; Brambilla *et al.*, 2010; Bräuer *et al.*, 2011; Yashiro *et al.*, 2011). It is known that when applying phylogenomics to

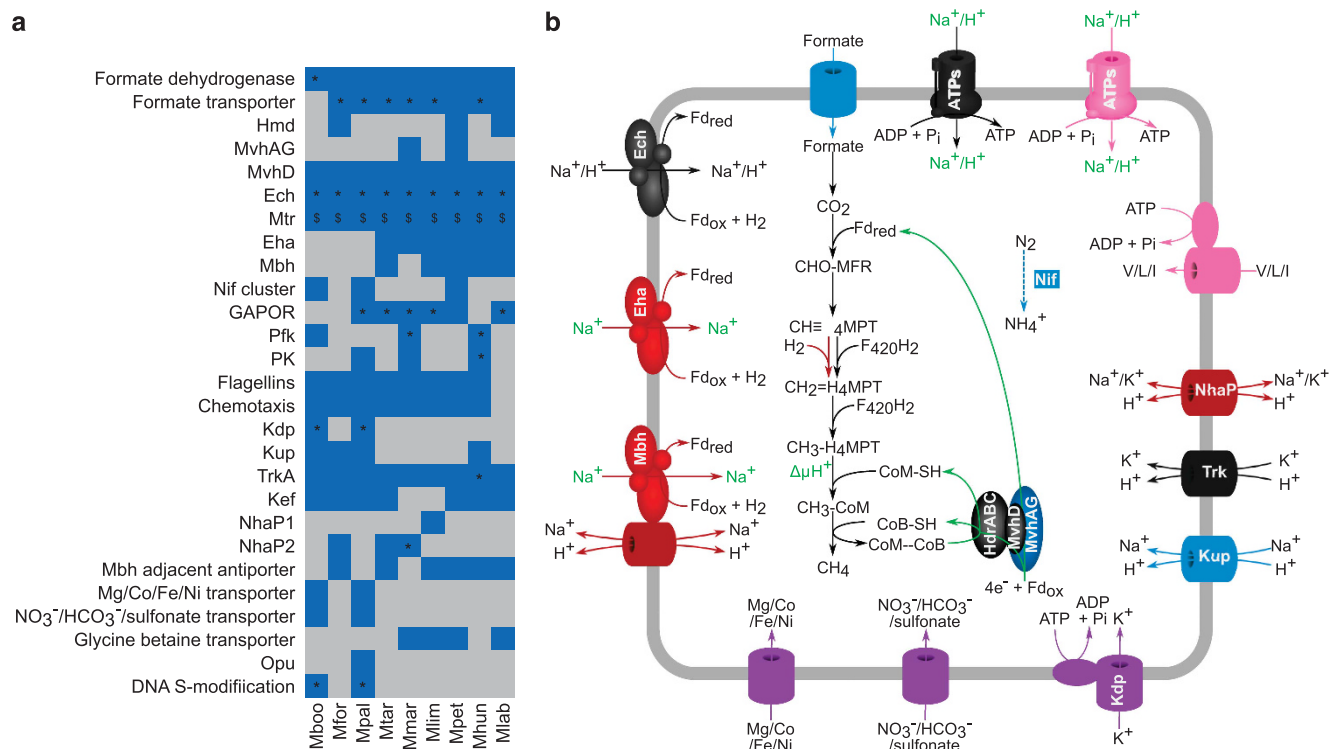


Figure 4 Adaptations of *Methanomicrobiales* to contrasting habitats. **(a)** Heatmap showing presence (blue) and absence (gray) patterns of various processes in the nine *Methanomicrobiales* isolates. ‘*’ denotes HGT and ‘s’ denotes positive selection evidence (in at least one subunit of multisubunit systems). **(b)** Cellular overview highlighting many of the processes evident in part **(a)**. Features in black are present in all strains. Those in blue are present in some, but not all, strains with no relation to habitat. Features in purple are present in only *M. boonei* and *M. palustris* and are potentially adaptive to peat dwelling. Features in cyan are found only in bioreactor-dwelling strains. Processes in green require biochemical and/or genetic evidence to determine the nature of the electron source or translocated ion. ATPs, ATP synthase; V/L/I, valine/leucine/isoleucine; CHO-MFR, formyl-methanofuran; $\text{CH}\equiv\text{H}_4\text{MPT}$, methenyl-tetrahydromethanopterin; $\text{CH}_2=\text{MPT}$, methylene- H_4MPT ; $\text{CH}_3-\text{H}_4\text{MPT}$, methyl-tetrahydromethanopterin; Hmd, H_2 -dependent methylene- H_4MPT dehydrogenase; Mtr, methyl-tetrahydromethanopterin S-methyltransferase; Mvh, F_{420} -nonreducing hydrogenase; Mvh, F_{420} -nonreducing hydrogenase; Ech, energy conserving hydrogenase; Eha, energy harvesting hydrogenase; Mbh, membrane-bound hydrogenase; Nif, nitrogen fixation; GAPOR, glyceraldehyde-3-phosphate ferredoxin oxidoreductase; Pfk, phosphofructokinase; PK, pyruvate kinase; Kdp, ABC-type K^+ transporter; Kup, K^+ uptake protein; TrkA, K^+ uptake protein; Kef, K^+ efflux transporter; NhaP, Na^+/H^+ antiporter; Opu, outer membrane porin; $\text{Fd}_{\text{red/ox}}$, reduced/oxidized ferredoxin; Pi, inorganic phosphate. Strain name abbreviations are defined in Figure 1.

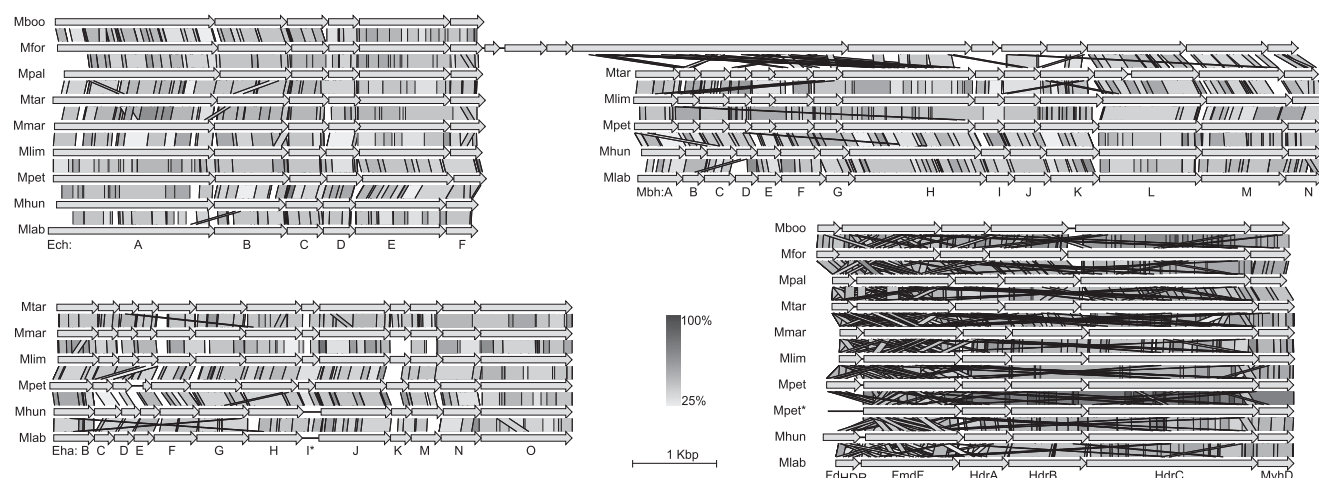


Figure 5 Synteny of membrane-bound hydrogenases and HdrED. Ech and Mbh subunits are not encoded adjacent to each other, except in the instance shown for *M. formicica*. **M. petrolearia* possesses two adjacent loci encoding the HdrED, with one of these loci lacking a ferredoxin encoding gene (FdHDR). The scale bars show the length corresponding to 1 kb and percent identity calculated from tblastx searches in grayscale. Abbreviations of strains are defined in Figure 1.

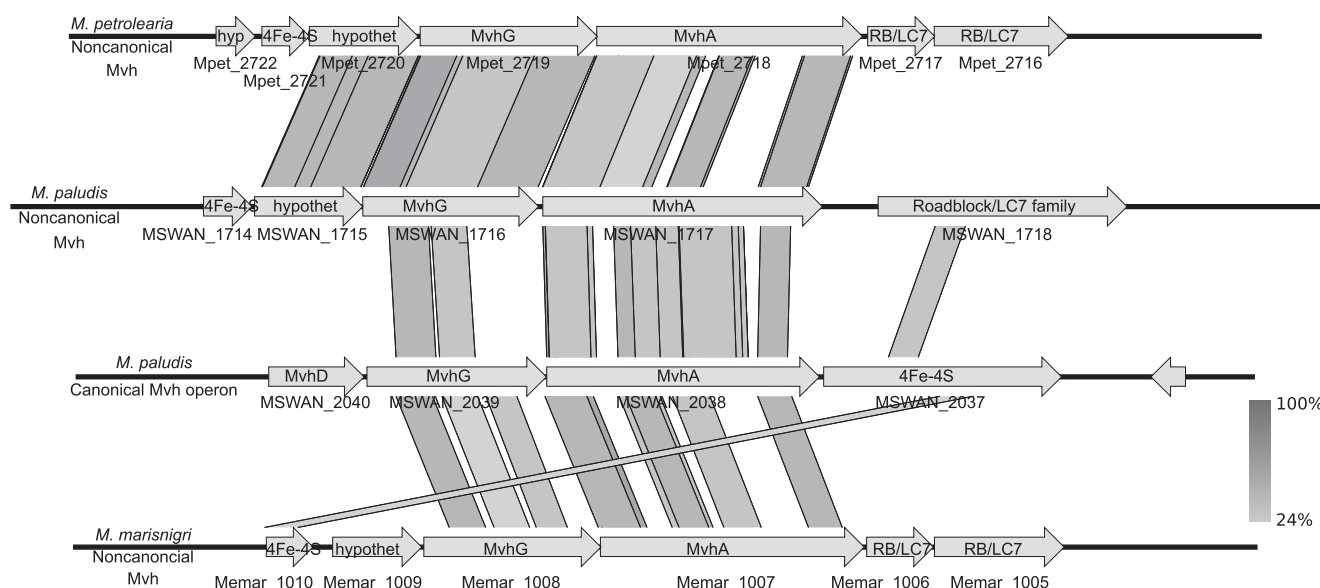


Figure 6 Synteny of F₄₂₀-nonreducing hydrogenases. The loci encoding Mvh subunits A and G from *M. marisnigri* and *M. petrolearia* are shown in comparison with canonical and non-canonical Mvh loci from *Methanobacterium paludis*. Locus tags are shown below the genes represented. Percent identities between loci were calculated using tblastx and are presented in grayscale. hyp, hypothetical protein; hypothet, hypothetical protein; RB/LC7, Roadblock/LC7 family protein.

distantly related organisms, the phylogenetic signal may become corrupted by long branch attraction artifacts (a particular issue for parsimony-based methods), through recombination and through substitution saturation (Philippe *et al.*, 2011). Our reconstructions indicate that *M. labreanum* and *M. hungatei* had long terminal branch lengths to nodes with relatively short internal branch lengths (Figure 1), indicating that the placement of these strains is not fully resolved yet and requires a few more related genomes to reduce this possible source of phylogenetic incongruence. HGT effects on core genome could be considered as an additional source of incongruence. However, concatenated approaches and Bayesian reconstruction, as done here, have been proposed as robust to low HGT frequencies (Lapierre *et al.*, 2014; Davidson *et al.*, 2015). Nevertheless, the incongruent branching order of the families between different data sets and methods indicates that future revisions of phylogenetic relationships between these families are required.

Gene flow dynamics evidence a different life history for *M. labreanum*

Assessment of expansion and/or streamlining of a genome can reveal putative adaptations (Ochman *et al.*, 2000; Lee and Marx, 2012). For instance, *M. labreanum* was the only strain in this study to evidence net gene loss since the divergence of the order. *M. labreanum* also evidenced differences in HGT, both in terms of acquisition rates (Supplementary Figure 4) and profile of HGT donor affiliations (Figure 3). These differences in *M. labreanum*'s HGT history cannot be accounted for by a lack of *Proteobacteria* in its habitat as *Proteobacteria* members are present in tar pits (Kim and Crowley, 2007). Notably,

the absence of archaellum and chemotaxis protein coding genes in *M. labreanum* is supported by phenotypic evidence showing the lack of motility or archaella under the microscope (Anderson *et al.*, 2009b). More generally, the genome of *M. labreanum* was deficient in genes associated with COG categories V (defense mechanisms) and T (signal transduction mechanisms) compared with the other strains (Supplementary Figure 5). This loss of secondary functions suggests genome downsizing leading to niche specialization (Lee and Marx, 2012). However, *M. labreanum*'s putative specialization is still unclear. For instance in other microbes, genome downsizing can sometimes indicate specialization to low substrate availability (Giovannoni *et al.*, 2005) but *M. labreanum*'s relatively rapid doubling time ($t_{1/2} = 12.5$ h) in pure culture indicates that it is capable of taking advantage of abundant substrate (H₂) supply.

Adaptations to peatlands

Methanoregula boonei and *M. palustris* were isolated as representatives of dominant methanogenic clades of contrasting peatlands, from an ombrotrophic bog (Bräuer *et al.*, 2006) and a minerotrophic fen (Cadillo-Quiroz *et al.*, 2008), respectively. Both strains must cope with low levels of inorganic nutrients and low pH (Dettling *et al.*, 2007), especially in the bog. Among the 60 clusters found only in peat-dwelling strains (Supplementary Table 2), an ATP-energized K⁺-transport system (Kdp) and two transport systems (one for cations and one for anions) of unknown specificity were evidenced. These three transport systems, along with a complete nitrogen fixation (*nif*) cluster (cluster Sδ, Supplementary Data 1) shared with *M. petrolearia*, suggest that adaptation to low availability of inorganic

nutrients is important for peat dwelling. Other protein clusters unique to the two peat-dwelling organisms include various metabolism-related proteins, putative heat-shock proteins/chaperones and a DNA sulfur modification system (possibly a defense-oriented host-specific restriction system; Xu *et al.*, 2010). Both peat-dwelling strains also appeared to show specializations in their iron uptake systems. While the other seven strains have gene clusters encoding several (two to nine) ABC-type iron transporters, *M. boonei* and *M. palustris* possess only one each. The fact that the only one type of substrate-binding protein is seen in the iron transporters of each peatland strain, while all other strains possess two to eight different substrate-binding domains in their iron transporters suggest that *M. boonei* and *M. palustris* have specialized iron acquisition systems.

Additionally, Kdp was identified as an HGT candidate of high importance for adaptation to peatland environments in *M. boonei* (Bräuer *et al.*, 2015; Cadillo-Quiroz *et al.*, 2015) and also in *M. palustris* (Bräuer *et al.*, 2015; Cadillo-Quiroz *et al.*, 2015). In peat soils, K⁺ concentrations are low (3–25 μ M) (Dettling *et al.*, 2007), but high levels of K⁺ are required intracellularly (up to 154 mM in *M. hungatei* GP 1; Sprott and Jarrell, 1981) for roles in pH homeostasis, cell turgor and membrane potential (Epstein, 2003). The two sets in *M. boonei* and one set of Kdp operons in *M. palustris* likely allow both organisms to overcome low K⁺ concentrations present in oligotrophic and minerotrophic peats. Kdp has a high affinity for K⁺ (K_m = 2 μ M in *Escherichia coli* K-12, with 37–63% protein identity to Kdp subunits of *M. boonei* and *M. palustris*) and has been shown to concentrate K⁺ in the cytoplasm when extracellular concentrations were as low as 50 nM (Ballal *et al.*, 2007). Our analyses of gene flow dynamics indicated that Kdp was likely acquired subsequent to the divergence of the *Methanoregula* family, in which the presence of Kdp operons in two peat-dwelling strains, *M. boonei* and *M. palustris*, and its absence in two bioreactor-dwelling ones, *Methanoregula formicica* and *M. tarda*, suggest that peatlands likely favor methanogens that acquired Kdp. As for the remaining strains in this study, all nine possess Trk- and some have Kup-type lower affinity K⁺-transport systems (Figure 4), both of which have K_m values around the order of 1 mM (Trchounian and Kobayashi, 1999; Epstein, 2003).

Our analyses indicated that the peatland methanogens, which are known to be highly sensitive to Na⁺ (Cadillo-Quiroz *et al.*, 2009; Bräuer *et al.*, 2011), are the only ones of this *Methanomicrobiales* study to not encode Na⁺-transporters. All other strains have a NhaP-type Na⁺/H⁺ antiporter and/or a multisubunit Na⁺/H⁺ antiporter adjacent to a membrane-bound hydrogenase (Figure 4). This corroborates the hypothesis that *M. boonei* has adapted to the low Na⁺/high H⁺ challenges of peat dwelling by using H⁺ rather than Na⁺ transmembrane chemiosmotic gradients (Bräuer *et al.*, 2015). Up to this point, based on studies of

Methanobacteriales and *Methanococcales* methanogens, it was thought that all cytochrome-deficient hydrogenotrophic methanogens conserve energy using a Na⁺ transmembrane chemiosmotic gradient, which is generated by tetrahydromethanopterin S-methyltransferase (Mtr) and consumed by a Na⁺-translocating ATP synthase (Schlegel and Müller, 2013). According to the new hypothesis, *Methanomicrobiales* members are expected to generate a H⁺ gradient using Mtr (as supported by previous analyses of the MtrE subunit; Bräuer *et al.*, 2015), and consume the H⁺ gradient, thus generating ATP, using a H⁺/Na⁺ promiscuous ATP synthase (Schlegel *et al.*, 2012; Bräuer *et al.*, 2015).

Adaptations to bioreactors and saline environments

The strains from bioreactors, *M. tarda*, *M. formicicum* and *M. hungatei*, all possessed another ATP synthase (cluster Se, in Supplementary Data 1) (in addition to the one core to the order), a branched-chain amino-acid transporter (cluster S ϕ), as well as a universal stress protein (cluster Sy) not seen in any of the other strains. *M. tarda* and *M. hungatei* shared yet another ATP synthase (cluster Sn) with *M. limicola*. Although all ATP synthases found in this order are predicted as Na⁺/H⁺ promiscuous type (Grüber *et al.*, 2014), alternative ATP synthases can have adaptive value for different growth rates or for different concentrations of Na⁺ and H⁺. The presence of a branched-chain amino-acid transporter indicates an ability to take advantage of substrates likely more available in eutrophic habitats, or environments with high microbial cell density and, possibly, significant dead microbial mass-turnover as in bioreactors.

All three *Methanomicrobiaceae* strains were isolated from saline environments (Romesser *et al.*, 1979; Wildgruber *et al.*, 1982; Ollivier *et al.*, 1997) and are also the most saline tolerant strains (0.69–0.86 M maximum NaCl tolerance) in this study. *M. labreanum* is also moderately NaCl tolerant (up to 0.51 M NaCl), while bioreactor strains are less tolerant to NaCl (up to 0.26 M NaCl). The peatland strains are the most NaCl sensitive of the strains in this study, tolerating no >0.1 M NaCl (Bräuer *et al.*, 2011). Microorganisms must maintain an osmotic pressure at least equivalent to their surroundings, which may be achieved by (i) the accumulation of salts, usually K⁺ and Cl[−] and known as the ‘salt-in strategy’ or (ii) by the accumulation of organic osmotic solutes (Oren, 2015). Since all strains possess various K⁺-transporters, regardless of their NaCl tolerances (Figure 4), it would seem that the accumulation of K⁺ is not necessarily the means by which *Methanomicrobiales* strains tolerate more or less NaCl. A putative glycine-betaine transporter was found in all three *Methanomicrobiaceae* strains and also in *M. labreanum* (cluster S ϕ) and not in any of the other strains (Figure 4). This organic solute transporter is proposed as a candidate for mediating NaCl/osmotic stress tolerance in the *Methanomicrobiales*. As the media in which the maximum NaCl tolerances were determined for the three *Methanomicrobiaceae* strains

and *M. labreanum* contained complex ingredients (for example, yeast extract and peptone hydrolysate) it is possible that these media contained suitable osmotic solutes for cytoplasmic osmolarity homeostasis. However, *de novo* synthesis of osmotically active solutes cannot be discounted.

It is important to consider that the genome availability of NaCl tolerant *Methanomicrobiales* is rather limited, and no genome sequences are currently available for strains within the most NaCl tolerant family of the order, the *Methanocalculaceae*, or for *Methanomicrobiaceae* strains isolated from wetlands or bioreactors (non-marine environments), hence it is likely that further genomic strategies to cope with NaCl/osmotic stress may yet become apparent within the *Methanomicrobiales*.

Analysis of primary metabolic genes reveals different lifestyles within the Methanomicrobiales

In hydrogenotrophic methanogens membrane-bound hydrogenases, such as Ech, Eha and Mbh, would be expected to catalyze the endergonic reduction of ferredoxin with H_2 at the expense of a transmembrane chemiosmotic gradient. Considering (i) our hypothesis that *M. boonei* and *M. palustris* utilize H^+ - rather than Na^+ -transmembrane chemiosmotic gradients, (ii) that *M. boonei* and *M. palustris* are the only strains in this study that appear to lack Na^+/H^+ antiporters and (iii) that Ech is the only membrane-bound hydrogenase apparent in the genomes of *M. boonei* and *M. palustris*, it could be expected that Ech is capable of translocating H^+ . Our results indicate that multi-subunit Na^+/H^+ antiporters are adjacent to Mbh (where present) and adjacent to Ech in *M. formicica* (Figure 5). This suggests that some of the *Methanomicrobiales* could concurrently use Na^+ and H^+ transmembrane chemiosmotic gradients as proposed in the *Methanosarcinales* (Schlegel and Müller, 2013). Ultimately, experimental investigations into the nature of ions translocated by the membrane-bound hydrogenases (Ech, Eha and Mbh), Mtr and ATP synthases will be required to understand the energy conservation and transduction mechanisms in the *Methanomicrobiales*. As a consequence of having only one membrane-bound hydrogenase, it would be expected that *M. boonei* and *M. palustris* are incapable of using separate membrane-bound hydrogenases to discriminate between anabolic and anaplerotic pools of ferredoxin as seen in *Methanococcus maripaludis* (Lie *et al.*, 2012), though they may possess another means of such discrimination.

In addition to differences in energy conservation pathways, these analyses also evidenced putative differences in the pathways of electron flow. Balancing cellular pools of reducing equivalents (H_2 , co-factor F_{420} ($F_{420}/F_{420}H_2$) and ferredoxin) is important as these reducing equivalents are required during several steps of methanogenesis and anabolic metabolism, and moreover, maintaining a suitably low H_2 concentration is important to the thermodynamic

constraints of syntrophic metabolism (Schink, 1997; Walker *et al.*, 2012). An F_{420} -reducing alcohol dehydrogenase (MMPG_2660) was found in *M. marisnigri*, *M. petroleariapetrolearius* and *M. tarda*. The former two are verified to be able to utilize secondary alcohols (Maestrojuán *et al.*, 1990; Ollivier *et al.*, 1997), while *M. tarda* was unable to utilize 2-propanol and 2-butanol (Imachi *et al.*, 2008). All strains had a cytoplasmic F_{420} -reducing hydrogenase, but they showed differential gene content at other steps of methanogenesis. In distantly related orders of cytochrome-deficient methanogens, the *Methanobacteriales* and *Methanococcales*, it was shown, during hydrogenotrophic methanogenesis, that two pairs of electrons are bifurcated in a process whereby the exergonic reduction of the coenzyme M-coenzyme B heterodisulfide is energetically coupled to the endergonic production of formyl-methanofuran, in a protein complex that incorporates cytoplasmic heterodisulfide reductase (HdrABC), formyl-methanofuran dehydrogenase, subunit D of Mvh and one of either formate dehydrogenase (Fdh), MvhABG, or an F_{420} -reducing hydrogenase (Costa *et al.*, 2010) (Figure 4). The proximity of MvhD to HdrABC (Figure 5) suggests that this subunit could interface with HdrABC during electron bifurcation as previously demonstrated in the *Methanococcales* order (Costa *et al.*, 2013). However, the organization of Mvh subunits (Figure 6) and lack of homologs to MvhB differs to what has been observed in *M. maripaludis* (Costa *et al.*, 2010) where experimental evidence showed that proximally encoded MvhABGD subunits (canonical enzyme system) mediated electron bifurcation. Overall, this indicates that there are profound differences in the mechanisms of electron bifurcation (particularly in the generation of low potential electrons) within the *Methanomicrobiales* and between the *Methanomicrobiales* and other orders of cytochrome-deficient methanogens. The implications of these differences in MvhABG content are not clear, as the pathways of electron bifurcation in hydrogenotrophic methanogens are still being elucidated and have been studied only in *Methanobacteriales* and *Methanococcales*. However, it may be of particular interest that Fdh increases in relative abundance in electron bifurcating complexes at the expense of MvhABG in *M. maripaludis* when hydrogen is in limiting supply (Costa *et al.*, 2013). This is interesting because Fdh is encoded in the *Methanomicrobiales'* core genome, including *M. boonei*, which is the only strain to lack a formate transporter and be unable to utilize formate as a methanogenic substrate (Bräuer *et al.*, 2011). This suggests a possible role for Fdh in electron bifurcation in the *Methanomicrobiales*, although experimental evidence is still pending.

The genome of *M. palustris* encodes a putative alternative membrane-bound HdrED, a key enzyme complex of the cytochrome containing *Methanosarcinales* (Buan and Metcalf, 2010). Homology search results indicate that *M. palustris* is the only archaeon outside the *Methanosarcinales* to possess a putative HdrE. HGT and maximum likelihood

analyses (Supplementary Figure 6) indicate that these genes could have been horizontally acquired from a *Methanosaeta* lineage, although more HdrED-containing genomes will improve this test. In terms of potential for functionality, there did not appear to be any means of synthesizing the methanophenazine electron carriers required by HdrED encoded in the genome of *M. palustris*, suggesting that HdrED may not have a canonical role in methanogenesis of *M. palustris*. However, the role and evolutionary history of HdrED, or HdrD alone, across methanogens is intriguing in light of this, plus new findings of HdrD presence (Borrel *et al.*, 2014) and active functioning (in the absence of HdrE) as HdrED for energy conservation in *Methanomassiliicoccus* (Kröninger *et al.*, 2016).

Two methylene-tetrahydromethanopterin (methylene- H_4 MPT) dehydrogenases, the F_{420} -dependent Mtd and the H_2 -dependent Hmd, catalyze the same step of methanogenesis, the reduction of methenyl- H_4 MPT to methylene- H_4 MPT. Expression studies have shown that one of Mtd or Hmd takes on quantitatively greater importance in response to H_2 limitation, rapid growth rate and syntrophic growth (Hendrickson *et al.*, 2007; Kato *et al.*, 2008; Enoki *et al.*, 2011; Walker *et al.*, 2012). In this study, Mtd was found in all strains, while Hmd was only found in three (Figure 4). This suggests that the adaptation to low H_2 concentrations afforded by Mtd is highly relevant for *Methanomicrobiales* under syntrophic lifestyles. Moreover, codon alignments of five F_{420} -dependent genes (Mtd, methylene- H_4 MPT reductase, and three out of four subunits of the cytoplasmic F_{420} -reducing hydrogenase) and three subunits of the energy conserving Mtr evidenced positive selection (Supplementary Data 2). This indicates that electron trafficking and energy conservation systems are subject to selection in these organisms. When also taking into account that many genes involved in central anabolic pathways such as acetate assimilation, amino-acid biosynthesis and gluconeogenesis are also subject to positive selection (Supplementary Data 2), it would appear that both energy metabolism and growth-related systems are under selective pressure. Such selective pressure can derive from syntrophic interactions, which can be subject to tight thermodynamic constraints. It is known that methanogens undergo gene regulation programs, impacting greatly on methanogenesis (energy production) and anabolic reactions (growth), to optimize syntrophic interactions (Shimoyama *et al.*, 2009; Walker *et al.*, 2012).

Conclusions

Here, we presented predicted differences in energy conservation, electron trafficking and other metabolic functions from the genomic make up of nine organisms within the *Methanomicrobiales*. Although different evidence lines converge in several of such putative predictions (Figure 4a), they require future examination via biochemical/genetic studies to

understand the physiology of *Methanomicrobiales* as distinct from other methanogens. The importance of this point is highlighted by the fact that seemingly subtle differences in energy conservation and electron trafficking pathways can be central to different lifestyles in contrasting environments, as seen, for example, in freshwater versus marine *Methanosarcina* spp. (Ferry and Lessner, 2008; Kulkarni *et al.*, 2009).

Comparative genomic analyses at the order level have allowed the proposition of a number of possible adaptations and differences across *Methanomicrobiales* strains from distinct taxonomic families and environments. The results of differential gene content, HGT, selection signatures and gene flow dynamics allow elaborating several hypotheses to further our understanding of the adaptations of methanogens to vastly different ecosystems.

Conflict of Interest

The authors declare no conflict of interest.

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